## GTCGACCCACGCGTCCGCGGACGCGTGGGCGGACGCGTGGGCGG

ACGCGTCCGGGGGACCGGTCGGGCCGGGACCAAGGGCACC	M S S	G A K	E G G	9
	ATG TCG TCC	GGG GCC AAG	GAG GGA GGT	27
G G S P A Y H L P H P GGG GGC TCT CCC GCC TAC CAC CTC CCT CAC CCC	H P H	P P Q	H A Q CAC GCC CAA	29 87
Y V G P Y R L E K T L TAT GTG GGC CCC TAT CGG CTG GAG AAG ACG CTG	g k g	Q T G	L V K	49
	GGC AAA GGA	CAG ACA GGG	CTG GTT AAA	147
L G V H C I T G Q K V CTC GGG GTC CAC TGC ATC ACG GGT CAG AAG GTC	A I K	I V N	R E K	69
	GCC ATC AAG	ATC GTG AAC	CGG GAG AAG	207
L S E S V L M K V E R CTG TCG GAG TCG GTG CTG ATG AAG GTG GAG CGG	E I A	I L K	L I E	89
	GAG ATC GCC	ATC CTG AAG	CTC ATC GAA	267
H P H V L K L H D V Y CAC CCA CAT GTC CTC AAG CTC CAC GAC GTC TAC	E N K	K Y L	Y L V	109
	GAG AAC AAG	AAA TAT TTG	TAC CTG GTT	327
L E H V S G G E L F D CTG GAG CAC GTC TCG GGG GGT GAG CTA TTC GAC	Y L V	K K G	R L T	129
	TAC CTG GTA	AAG AAG GGG	AGA CTG ACG	387
P K E A R K F F R Q I CCC AAG GAG GCC CGA AAG TTC TTC CGC CAG ATT	V S A	L D F	C H S	149
	GTG TCT GCG	CTG GAC TTC	TGC CAC AGC	447
Y S I C H R D L K P E TAC TCC ATC TGC CAC AGA GAC CTA AAG CCC GAG	N L L	L D E	K N N	169
	AAC CTG CTT	TTG GAT GAG	AAA AAC AAC	507
I R I A D F G M A S L	Q V G	D S L	L E T	189
ATC CGC ATT GCA GAC TTC GGC ATG GCG TCC CTG	CAG GTG GGG	GAC AGC CTC	CTG GAG ACC	567
S C G S P H Y A C P E AGC TGC GGG TCC CCC CAT TAT GCG TGT CCA GAG	V I K	G E K	Y D G	209
	GTG ATT AAG	GGG GAA AAA	TAT GAT GGC	627
R R A D M W S C G V I	L F A	L L V	G A L	229
CGC CGG GCA GAC ATG TGG AGC TGT GGA GTC ATC	CTC TTC GCC	CTG CTC GTG	GGG GCT CTG	687
P F D D D N L R Q L L CCC TTT GAT GAC GAC AAC CTC CGC CAG CTG CTG	E K V	K R G	V F H	249
	GAG AAG GTG	AAA CGG GGC	GTC TTC CAC	747
M P H F I P P D C Q S	L L R	G M I	E V E	269
ATG CCC CAC TTC ATT CCT CCA GAT TGC CAG AGC	CTC CTG AGG	GGA ATG ATC	GAA GTG GAG	807
P E K R L S L E Q I Q CCC GAA AAA AGG CTC AGT CTG GAG CAA ATT CAG	K H P	W Y L	G G K	289
	AAA CAT CCT	TGG TAC CTA	GGC GGG AAA	867
H E P D P C L E P A P CAC GAG CCA GAC CCT	G R R	V A M	R S L	309
	GGC CGC CGG	GTA GCC ATG	CGG AGC CTG	927
P S N G E L D P D V L	E S M	A S L	G C F	329
CCA TCC AAC GGA GAG CTG GAC CCC GAC GTC CTA	GAG AGC ATG	GCA TCA CTG	GGC TGC TTC	987
R D R E R L H R E L R	S E E	E N Q	E K M	349
AGG GAC CGC GAG AGG CTG CAT CGC GAG CTG CGC	AGT GAG GAG	GAG AAC CAA	GAA AAG ATG	1047
I Y Y L L L D R K E R	Y P S		Q D L	369
ATA TAT TAT CTG CTT TTG GAT CGG AAG GAG CGG	FAT CCC AGC		CAG GAC CTG	1107
P P R N D V D P P R K CCT CCC CGG AAT GAT GTT GAC CCC CCC CGG AAG		S P M TCT CCC ATG		389 1167
CAC GGG AAG CGG CGA CCA GAG CGG AAG TCC ATG	GAA GTC CTG		GAT GCC GGG	409 1227
G G G S P V P T R R A GGT GGT GGC TCC CCT GTA CCC ACC CGA CGG GCC	TTG GAG ATG	GCC CAG CAC	AGC CAG AGA	429 1287
S R S V S G A S T G L TCC CGT AGC GTC AGT GGA GCC TCC ACG GGT CTG				449 1347

Fig. 1B

## Analysis of 55053 (778 aa)

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>55053 MSSGAKEGGGGSPAYHLPHPHPPPQHAQYVGPYRLEKTLGKGQTGLVKLGVHCITGQKV AIKIVNREKLSESVLMKVEREIAILKLIEHPHVLKLHDVYENKKYLYLVLEHVSGGELFD YLVKKGRLTPKEARKFFRQIVSALDFCHSYSICHRDLKPENLLLDEKNNIRIADFGMASL QVGDSLLETSCGSPHYACPEVIKGEKYDGRRADMMSCGVILFALLVGALPFDDDVLRQLL EXVKRGVFHMPHFIPPDCQSLLRGMIEVEPEKRLSLEQIQKHPWYLGGKHEPDPCLEPAP GRRVAMRSLPSNGELDPDVLESMASLGCFRDRERLHRELRSEEENQEKMIYYLLLDRKER YPSCEDQDLPPRNDVDPPRKRVDSPMLSRHGKRRPERKSMEVLSITDAGGGGSPVPTRRA LEMAQHSQRSRSVSGASTGLSSSPLSSPRSPVFSFSPEPGAGDEARGGGSPTSKTQTLPS RGPRGGGAGEQPPPPSARSTPLFGPFGSPRSSGGTPLHSPLHTPRASPTGTPFPFSP SLDKEEQIFLVLKDKPLSSIKADIVHAFLSIPSLSHSVLSQTSFRAEYKASGGPSVFQKP VRFQVDISSSEGPEPSPRRDGSGGGGIYSVTFTLISGPSRRFKRVVETIQAQLLSTHDQP SVQALADEKNGAQTRPAGAPPRSLQPPPGRPDPELSSSPRRGPFKDKKLLATNGTPLP **GGGVGGAAWRSRLNSIRNSFLGSPRFHRRKMQVPTAEEMSSLTPESSPELAKRSWFGNFI** 

## Transmembrane Segments Predicted by MEMSAT

Score	4.1	9.0	1.2
Orient	ins>out	out>ins	ins>out
End	231	640	697
Start	214	624	188

## Protein Family / Domain Matches, HMMer version 2

```
Searching for complete domains in PFAM
hmmpfam - search a single seq against HMM database
HMMER 2.1.1 (Dec 1998)
Copyright (C) 1992-1998 Washington University School of Medicine
HMMER is freely distributed under the GNU General Public License (GPL).
                         /prod/ddm/seqanal/PFAM/pfam5 5/Pfam
                         /prod/ddm/wspace/orfanal/oa-script.23506 seq
Sequence file
 Query: 55053
Scores for sequence family classification (score includes all domains)
                                                       Score E-value N
Model Description
                                                                 2.6e-93
pkinase Eukaryotic protein kinase domain
                                                                     4.9
         UBA domain
JBA
Parsed for domains
                                                score E-value
Model Domain seq-f seq-t
                             hmm-f hmm-t
                        285 .. 1 278 []
                                                323.4 2.6e-93
           1/1
                   34
pkinase
                                                  7.7
                        356 ..
                                   1
                                      41 ()
          1/1
Alignments of top-scoring domains:
pkinase: domain 1 of 1, from 34 to 285. score 323 4, E = 2.6e-93
                   •->yelleklGeGsfGkVykakhk.tgkıvAvKılkkesls.....lr
                      y+1 ++1G+G G V++++h tg++vA+K1+++e+1s++ + r
                      YRLEKTLGKGQTGLVKLGVHC1TGQKVAIKIVNREKLSesvlmkvER 80
       55053
                   {	t ElqilkrlsHpNIvrllgvfedtddhlylvmEymegGdLfdylrrngpls}
                   E1+1lk + Hp++++l++v+e +++lylv+E++ gG+Lfdyl+++g+l+
                81 EIAILKLIEHPHVLKLHDVYE-NKKYLYLVLEHVSGGELFDYLVKKGRLT 129
       55053
                   ekeakkıalQılrGleYLHsngıvHRDLKpeNILldengtvKiaDFGLAr
                   +kea+k+++Q1+++l+++Hs +1+HRDLKpeN+Llde+++++iaDFG+A
               130 PKEARKFFRQIVSALDFCHSYSICHRDLKPENLLLDEKNNIRIADFGMAS 179
       55053
                   ll...eklttfvGTpwYmmAPEvileg.rgysskvDvWSlGviLyElltg
                      +++ 1 t +G+p+Y PEV ++g+++++++D+WS+GviL+ 11 g
               180 LQvgdSLLETSCGSPHYA-CPEV-IKGeKYDGRRADMWSCGVILFALLVG 227
       55053
                   qplfpqadlpaftggdevdqliifvlklPfsdelpktridpleelfrikk
                                                          d+1++1++ +k
                                              lPf+d
               228 -----DNLRQLLEKVK 244
       55053
                   r.rlplpsncSeelkdLlkkcLnkDPskRpGsatakeilnhpwf<-*
               r+ + p+ ++++++Ll+++++ P+kR+ + ++i +hpw
245 RgVFHMPHFIPPDCQSLLRGMIEVEPEKRL---SLEQIQKHPWY
 UBA: domain 1 of 1, from 315 to 356: score 7.7, E = 4.9
                     ->edeekieqLveMGF..dreevvkALratngngverAaewLlsh<-*
                       d + +e++ ++G +dre+ + Lr+ n e+ +++Ll +
                      LDPDVLESMASLGCfrDRERLHRELRSEEEN-QEKMIYYLLLD
        55053
               315
 Searching for complete domains in SMART
 hmmpfam - search a single seq against HMM database
 HMMER 2.1.1 (Dec 1998)
 Copyright (C) 1992-1998 Washington University School of Medicine
 HMMMER is freely distributed under the GNU General Public License (GPL).
                          /ddm/robison/smart/smart.all.hmms
 HMM file:
                          /prod/ddm/wspace/orfanal/oa-script.23506.seq
 Sequence file:
  Query: 55053
```

Fig. 3A

Scores for sequence family classification (score includes all domains): E-value N Model Description Score 2.4e-103 1 -----356.8 serkın\_6 2 4e-14 1 39.2 tyrkın\_6 Parsed for domains Model Domain seg f seg-t hmm-f hmm t score E-value 1 231 () 1 280 () 34 285 34 286 serkin\_6 1/1 tyrkin\_6 1/1 356 8 2 4e-103 39 2 2 4e-14 Alignments of top-scoring domains serkin\_6: domain 1 of 1, from 34 to 285. score 356 8, E = 2 4e-103 \*->YellkklGkGaFGkVylardkktgrlvAlKvık .... erilr Y+l k+lGkG G V+l+++ tg++vAlK+++++ +++ +++r YRLEKTLGKGQTGLVKLGVHCITGQKVAIKIVNreklsesvlMKVER 80 55053  ${\tt EikiLkk~dHPNIVkLydvfed~dklylVmEyceGdlGdLfdllkkrgrr}$ E1+1Lk HP++ kL+dv+e++++lylv+E+++G G+Lfd+l+k+gr 81 EIAILKL1EHPHVLKLHDVYENKKYLYLVLEHVSG--GELFDYLVKKGR- 127 55053 glrkvlsE.earfyfrQilsaLeYLHsqgIiHRDLKPeNiLLds .hvKl l+++ear++frQ1+saL+++Hs I+HRDLKPeN+LLd+++++ 128 ----LTPKEARKFFRQIVSALDFCHSYSICHRDLKPENLLLDEKNNIRI 172 55053 aDFGlarql....ttfvGTpeYmAPEvl..gYgkpavDiWSlGcily aDFG+A + +++ t +G+p+Y PEv++++Y++++D+WS+G+il+ 173 ADFGMASLQvgdsllETSCGSPHYACPEVIkgeKYDGRRADMWSCGVILF 222 55053 ElltGkpPFp..qldlifkkig...... SpeakdLıkklLvkdPek ll+G PF++++l ++++k++++ + ++ +p++ +L++++ +++Pek 223 ALLVGALPFDddNLRQLLEKVKrgvfhmphf1PPDCQSLLRGMIEVEPEK 272 55053 Rlta.eaLedeldikaHPff<-\* R1+ +++ + HP+ 273 RLSLeQIQK-----HPWY 55053 285

Fig. 3B

tyrkin_6: domain 1	of 1, from 34 to 286: score 39.2, E = 2.4e-14 *->ltlgkkLGeGaFGeVykGtlkieVAVKtLkedakeeFlr	
55053 34	+1+k+LG G+ G V +G+ ++++VA+K ++ ++ +++ r YRLEKTLGKGQTGLVKLGVHCitgQKVAIKIVNREK1sesvLMKVER	80
	Eak1MkklGgkHpNiVkLlGvcteegrrFmevePlmivmEymeqGdLldy	
	E+ i+k + +Hp+++kL+ v + + 1++v+E+++gG L dy	
55053 81	EIAILKLIEHPHVLKLHDVYENKKYLYLVLEHVSGGELFDY	
	LrknrpklslsdLlsfAlQIAkGMeYLesknfvHRDLAARNcLvgenkvv	
	L k+++ 1++++ +f QI + ++ +s + HRDL N L++e++ +	
55053 122	LVKKGR-LTPKEARKFFRQIVSALDFCHSYSICHRDLKPENLLLDEKNNI	170
	${\tt KIsDFGLsRdlyddDkkGeskdyYrkkggkggktllPirWmAPEslkdgk}$	
	+I+DFG++ d $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$	
55053 171	RIADFGMASLQVGDSLLETSCGSPHYACPEVIKGEK	206
	${\tt Ft.skSDVWSFGVlLWE} if {\tt tlGeqPYpgelqqfmsneevleylkkGyRlp}$	
EE052 207	+ + D WS GV L+ ++ G+ P + ++++le++k+G	
55053 207	YDgRRADMWSCGVILFALL-VGALPFDDDNLRQLLEKVKRG-VFH	249
	kPendlpiSsvtCPdelYdlMlqCWaedPedRPtFselverl<-* P+ P++++++++++++++++++++++++++++++++++	
55053 250		
55053 250	MPHFIPPDCQSLLRGMIEVEPEKRLSLEQIqkhPWYL 28	56
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Fig. 3C